(1) Publication number:

0 184 309

(12)

EUROPEAN PATENT APPLICATION

(21) Application number: 85307789.9

(5) Int. Cl.4: **C 07 K 7/06** C 07 K 7/08, A 61 K 37/02

22 Date of filing: 28.10.85

30 Priority: 01.11.84 GB 8427651

43) Date of publication of application: 11.06.86 Bulletin 86/24

(84) Designated Contracting States: BE CH DE FR GB IT LI LU NL SE 71 Applicant: BEECHAM GROUP PLC **Beecham House Great West Road Brentford Middlesex TW8 9BD(GB)**

(72) Inventor: Summers, Christine The Patch The Street Sturmer Essex, CB9 7XF(GB)

nventor: Wootton, Gordon 51 Parkway Sawbridgeworth Hertfordshire(GB)

(72) Inventor: Watts, Eric Alfred 217 Waterhouse Moor Harlow Essex, CM18 6BW(GB)

(74) Representative: Jones, Pauline et al, Beecham Pharmaceuticals Patents & Trade Marks Dept. **Great Burgh Yew Tree Bottom Road** Epsom Surrey KT18 5XQ(GB)

[54] Peptides selected from amino-acid residues 11 to 23 of VIP.

(57) Peptides comprising, in sequence, units selected from the amino acid residues 11 to 23 of vasoactive intestinal peptide (VIP) and consisting at least of the amino acid residues 15 to 20, or an analogue thereof wherein one or more of the amino acid residues is replaced by an equivalent other amino acid, or a pharmaceutically acceptable salt thereof; having pharmacological activity, a process for their preparation and their use as pharmaceuticals.

. 1 -

02.

Novel Compounds

The invention relates to VIP fragments and analogues, processes for their preparation, pharmaceutical preparations containing them and their use in medicine.

Vasoactive intestinal peptide (VIP) was originally isolated from the small intestines of the hog, but it has since been isolated from other species, such as the chicken, and has been shown to have a wide distribution throughout body tissues.

It has systemic vasodilator activity. It induces systemic hypotension and increases cardiac output on intravenous infusion. It increases hepatic arterial blood flow, increases blood sugar levels, and has the ability to bring about tracheal relaxation, and relaxation of gut smooth muscle, as well as stimulation of the output of bicarbonate from intestinal secretions. It therefore appears to be useful in treatment of hypertension and peripheral vascular disease on parenteral administration, and as a bronchodilator on aerosol or parenteral administration.

Vasoactive intestinal peptide comprises a peptide having a sequence of 28 amino acids in a single chain. The sequence of VIP (pig) is shown in table 1.

Table 1
VIP (pig)

N-Terminus

N—NH CH ₂ H ₂ NCHCO	он . Сн ₂ .	CO ₂ H CH ₂ I	— NECHCO — CH3	CH CH3	CH ₂ —NHCHCO —	CH OH
His	Ser	Asp	Ala	Val	Phe	Thr
СО ₂ Н СН ₂ — NHCHCO —	CONH 2 CH 2 CH 2 NHCHCO —	OH CH ₂ -NHCHCO	CH CH NHCHCO —	C NH C NH C NH C C S S C C C C C C	CH ₃ CH ₃ CH CH ₂ NHCHCO	H ₂ N NH C I NH (CH ₂) 3
Asp	Asn	Tyr	Thr	Arg	Leu	Arg
NH2 CH2)4 -NHCHCO-	CONH 2 (CH 2) 2 NHCHCO —	CH ₃ s (CH ₂) ₂ -NHCHCO	СН _З — МНСНСО — Ala	CH 3 CH 3 CH 1 NHCHCO—	NH 2 2 (CH 2) 4 	NH ₂ (CH ₂) ₄ NHCHCO—
CH ₂ -NHCHCO-	CH 3 CH 3 CH 1 CH 2 NHCHCO —	CONH ₂ CH ₂ NHCHCO -	OH CH2 NHCHCO — Ser	CH ₃ CH ₂ CH ₃ CH NHCHCO	CH ₃ CH ₃ CH CH ₂ NHCHCO	CONH 2 CH 2 NHCHCONH 2 Asn-NH 2

C-Terminus

Abbreviations used are as follows:

υ	J
	_

04	Amino Acid Residue	Abbreviations
05		
06	alanine	Ala
07	arginine	Arg
08	asparagine	Asn
09	aspartic acid	Asp
10	glutamine	Gln
11	histidine	His
12	isoleucine	Ile
13	leucine	Leu
14	lysine	Lys
15	methionine	Met
16	norleucine	Nle
17	phenylalanine	Phe
18	serine	Ser
19	threonine	Thr
20	tyrosine	Tyr
21	valine	Val

The amino acid components are in the L-form.

VIP (chicken) is closely related, differing in the 11, 13, 26 and 28 positions; the peptide has:

a serine residue in position 11, a phenylalanine residue in position 13, a valine residue in position 26 and a threonine residue in position 28.

A number of C-terminal fragments have been produced, mostly in the synthetic programme required to prove the structure of VIP. Few structures have been obtained from the N-terminus, and very little work has been

01	4 -
02	carried out on fragments from the centre of the
03	molecule.
04	
05	It has, however, been concluded (Robberecht, Gut
06	Hormones (1978) edited by Bloom, p 97 to 103) that the
07	C-terminus of VIP holds the receptor recognition site,
08	and that the N-terminus holds the activation site,
09	together with a minimal capacity for binding.
10	
11	Counter to the commonly held views regarding the
12	activity of VIP, we have found that there is
13	pharmacological activity even in the absence of the
14	amino acid units at the C- and N-termini of the
15	molecule.
16	-
17	The present invention provides a peptide comprising, in
18	sequence, units selected from the amino acid residues
19	11 to 23 of VIP and consisting at least of the amino
20	acid residues 15 to 20, or an analogue thereof wherein
21	one or more of the amino acid residues is replaced by
22	an equivalent other amino acid.
23	
24	The present invention also provides a peptide
25	consisting, in sequence, of the VIP units selected from
26	the amino acid residues 11 to 23, and comprising at
27	least the amino acid residues 15 to 20, or an analogue

thereof having pharmacological activity.

28. 29 30

31

32

33

34

Preferably in a peptide of the present invention the amino acid units are selected from residues 13 to 23 or 11 to 21, more especially from residues 13 to 21, of In an analogue thereof, one or more than one amino acid unit may be replaced by an equivalent amino acid unit.

- 5 -

Amino acids can be considered as members of different classes; such groupings are well known. Replacement of an amino acid of the peptide by an equivalent amino acid may be by another amino acid of the same class, and where an amino acid can be grouped into two or more classes, replacement may be made from one or more of these classes.

All amino acids in an analogue of the present invention may, for example, be naturally occurring amino acids, i.e. L-amino acids, or amino acids in the D- or DL-form.

 It seems reasonable to suppose that the activity of a pertide bears some relationship to its secondary structure (which could be inherent, or adopted at the receptor site). Thus the expressed activity could be related to a potential for formation of a highly ordered arrangement of some of the amino acids.

Where there is replacement of one or more amino acids, the replacement may, for example, be such that the essential structure of the fragment is maintained.

Without intending to be limited to the following hypothesis, we presently believe it is possible that, for peptides of the present invention, a helical structure may be a contributory factor in the pharmacological activity. The replacing amino acid or acids in an analogue thereof may therefore, if desired, be selected so as to have at least as good a helical-forming character as the replaced amino acid(s). However, lack of a helical structure may not impair the activity of a peptide or analogue of the present invention; for example, it may be preferred, for pharmacological reasons or otherwise, to

01	- 6 -
02	incorporate D-amino acid(s) as the replacing amino
03	acid(s) and it will, of course, be understood that
04	unless all amino acids in the resulting analogue are in
05	the D-form, the structure will not be of a helical
06	nature.
07	
08	Thus, for example:
.)9	·
10	- the threonine at position 11 of VIP (pig) may, if
11	desired, be replaced by another hydroxy amino acid,
12	e.g. serine (Ser); and the serine at position 11 of VIP
13	(chicken) may, if desired, be replaced by another
14	hydroxy amino acid, e.g. threonine (Thr);
15	
16	- the arginine at position 12 and/or at position 14
17	may, if desired, be replaced by another basic amino
18	acid, e.g. lysine (Lys) or ornithine (Orn);
19	·
20	- the leucine at position 13 of VIP (pig) and/or at
21	position 23, and the phenylalanine at position 13 of
22	VIP (chicken) may, if desired, be replaced by another
23	hydrophobic amino acid, in the case of leucine, by, for
24	example, valine (Val), and, in the case of
25	phenylalanine, by, for example, tyrosine;
26	
27	- the lysine at any one or more of positions 15, 20 and
. 28,	21 may, if desired, be replaced by another basic amino
29	acid, e.g. ornithine (Orn) or arginine (Arg);
30	
31	- the glutamine at position 16 may, if desired, be
32	replaced by another carboxamido amino acid, e.g.
33	asparagine (Asn);
34	
35	- the methionine at position 17 may, if desired, be
36	replaced by another neutral amino acid, e.g. the iso-
37	steric norleucine (Nle) or leucine (Leu);
30	

01	- 7 -
0.2	- the alanine at position 18 may, if desired, be
03	replaced by another hydrophobic amino acid, e.g.
04	glycine (Gly) or norvaline (Nva);
05	
06	- the valine at position 19 may, if desired, be
07	replaced by another hydrophobic amino acid, e.g.
08	leucine (Leu);
09	
10	- the tyrosine at position 22 may, if desired, be
11	replaced by another hydrophobic amino acid, especially
12	an aromatic amino acid, e.g. phenylalanine (Phe).
13	
14	Especially, there should be mentioned analogues in
15	which one or more of the amino acid residues 15 to 20
16	is replaced by an equivalent other amino acid and any
17	additional amino acid residues present correspond to
18	those in VIP.
19	
20	Especially, the present invention provides a
21	hexapeptide amide with the amino acid sequences of the
22	residue 15 to 20 of VIP, or an analogue thereof in
23	which one or more of the amino acids is replaced as
24	indicated above.
²⁵	
26	Very especially, the present invention provides the
27	hexapeptide
28	
29	Lys Gln Y Ala Val Lys
30	•
31	where Y represents Met or Nle; and also the hexapeptide
32	
33	Lys Gln Y Ala Leu Lys
34	
35	where Y represents Met or Nle.
36	

CI	- 8 -
02	Fragments and analogues of VIP (pig) should especially
03	be mentioned, but the basic structure may correspond to
04	VIP from any source.
05	
06	The following fragments and analogues should especially
07	be mentioned:
08	
09	Arg [A];
10	Leu Arg [A];
11	Arg Leu Arg [A];
12	Thr Arg Leu Arg [A];
13	[A] Lys;
14	[A] Lys Tyr;
15	[A] Lys Tyr Leu;
16	Arg [A] Lys;
17	Arg [A] Lys Tyr;
18	Arg [A] Lys Tyr Leu;
19	Leu Arg [A] Lys;
20	Leu Arg [A] Lys Tyr;
21	Leu Arg [A] Lys Tyr Leu;
22	Arg Leu Arg [A] Lys;
23	Arg Leu Arg [A] Lys Tyr;
24	Arg Leu Arg [A] Lys Tyr Leu;
25	Thr Arg Leu Arg [A] Lys;
26	Thr Arg Leu Arg [A] Lys Tyr;
27	Thr Arg Leu Arg [A] Lys Tyr Leu.
28	
29	where
30	·
31	[A] denotes Lys Gln Y Ala Val Lys
32	·
33	in which Y represents Met or Nle.
34	
35	The amino acids may, for example, be in the L-form;
36	although one or more D-amino acids may, if desired, be
37	present in the structure.
38	

The carboxy-terminus of the peptides or analogues of the present invention may be in the form of the acid (-OH); an ester, for example an alkyl ester, especially a (C1-C4)-alkyl ester, e.g. the methyl ester, (-OCH3), the hydrazide (-NH-NH2), or an amide, usually the unsubstituted amide (-NH2). Preferably the carboxy-terminus is in the form of the unsubstituted amide.

The amino-terminus of the peptides or analogues of the present invention may be in the form of the unsubstituted amine (-NH₂) or protected amine (-NHR) where R represents, for example, acetyl, tert.-butyloxycarbonyl or benzyloxycarbonyl, or in the form of an acid addition salt, preferably a physiologically tolerable, pharmaceutically acceptable acid addition salt, of the amine.

Acid addition salts may be, for example, salts with inorganic acids such, for example, as hydrochloric acid, hydrobromic acid, orthophosphoric acid or sulphuric acid, or organic acids such, for example, as methanesulphonic acid, toluenesulphonic acid, acetic acid, trifluoroacetic acid, propionic acid, lactic acid, citric acid, tartaric acid, fumaric acid, malic acid, succinic acid, salicylic acid or acetylsalicylic acid.

Thus, more particularly, the present invention provides a polypeptide of the general formula

$$X - (Y')_n - Y_{15}Y_{16}Y_{17}Y_{18}Y_{19}Y_{20} - (Y'')_m - Z$$
 I

in which

X represents a hydrogen atom or an amine-protecting group, preferably a hydrogen atom;

02	$(Y^*)_n$ represents a direct bond or
03	
04	$Y_{11}Y_{12}Y_{13}Y_{14}$, $Y_{12}Y_{13}Y_{14}$, $Y_{13}Y_{14}$ or Y_{14}
05	
06	in which
07	
08	Y11 represents Thr, Ser or the residue of another
09	hydroxy amino acid,
10	
11	${ m Y}_{12}$ represents Arg or the residue of another basic
12	amino acid,
13	
14	Y13 represents Leu, Phe or the residue of another
15	hydrophobic amino acid,
16	
17	Y_{14} represents Arg or the residue of another basic
18	amino acid;
19	
20	Y_{15} represents Lys or the residue of another basic
21	amino acid, e.g. Orn,
22	
23	Y_{16} represents Gln or the residue of another carbox-
24	amido amino acid,
25 .	<u>.</u>
26	Y17 represents Met or the residue of another neutral
27	amino acid, e.g. Nle,
28	
29 .	Y_{18} represents Ala or the residue of another
30	hydrophobic amino acid,
31	
32	Ylg represents Val or the residue of another hydro-
33	phobic amino acid,
34	
35	Y20 represents Lys or the residue of another basic
. 36	amino acid, e.g. Orn,
37	

(Y'')_m represents a direct bond or

04 Y₂₁, Y₂₁Y₂₂ or Y₂₁Y₂₂Y₂₃

06 in which

Y21 represents Lys or the residue of another basic amino acid,

Y22 represents Tyr or the residue of another hydrophobic amino acid,

 Y_{23} represents Leu or the residue of another hydrophic amino acid; and

Z represents a hydroxyl group, or a group of the formula OR such that CÔZ represents an ester, or a hydrazino group such that COZ represents a hydrazide, or NH₂ such that COZ represents an amide, preferably NH₂; and salts thereof, preferably physiologically tolerable salts thereof, especially physiologically tolerable acid addition salts thereof.

The compounds of formula I are preferably in pharmaceutically acceptable form. By pharmaceutically acceptable form is meant, inter alia, of a pharmaceutically acceptable level of purity excluding normal pharmaceutical additives such as diluents

 carriers, and including no material considered toxic at normal dosage levels. A pharmaceutically acceptable level of purity will generally be at least 50% excluding normal pharmaceutical additives, preferably 75%, more preferably 90% and still more preferably 95%.

synthesis or solid phase procedures may be used.

The coupling reactions may be effected by, for example, activating the reacting carboxyl group of the ingoing amino acid, and reacting this with the amino group of the substrate unit. Details of suitable, optional activating and protecting (masking) groups and of suitable reaction conditions (for the coupling reactions and for the introduction and removal of protecting groups) giving, preferably, the minimum of racemisation, may be found in the above-referenced literature.

Accordingly, the present invention further provides a process for the preparation of a peptide or analogue of the present invention, which comprises coupling a suitable amino acid or amino acid sequence in which the carboxyl group is activated with an appropriate amino acid or amino acid sequence and repeating, if necessary, the coupling procedure until there is obtained a peptide comprising, in sequence, units selected from the amino acid residues 11 to 23 of VIP consisting at least of the amino acid residues 15 to 20, or an analogue thereof in which one or more of the amino acid residues is replaced by an equivalent other amino acid, wherein, if desired or required, non-reacting functional groups are protected during the coupling procedure and, if desired, subsequently deprotected.

A polypeptide of the general formula I may thus be prepared by reacting a reagent of the general formula

$$H - A_J - OH$$
 (II)

wherein

0 :	
02	Yl represents an amino acid unit or a partial radical
03	sequence identical with the corresponding N-terminal
04	amino acid unit or partial radical sequence in formula
05	I,
06	
07	with a reagent of the general formula
08	.
09	$H - Y^2 - OH$ (III)
10	
11	wherein
12	
13	Y ² represents an amino acid unit or a partial radical
14	sequence identical with that in the balance of the
15	above-defined product peptide,
16 .	
17	the reagents (II) and (III) being optionally protected
18	and/or activated where and as appropriate, followed if
19	desired or required by one or more of the following:
20	· -
21	- deprotection of the products,
22	
23	 conversion of one carboxy terminus into another
24	carboxy terminus,
25	
26	 conversion of a free peptide into a salt thereof.
27	
28	For example, an appropriate peptide ester of the
29	general formula
30	
31	$X - Y^1 - Y^2 - OR \tag{IV}$
32	
33 ·	wherein X, Y^1 and Y^2 have the meanings given above and
34	R represents, for example, an alkyl group and
35	preferably an alkyl group having 1 to 4 carbon atoms,
36	may be converted into an amide by reaction with
37	ammonia.

	U1043
01	- 15 -
02	Compounds of the general formulae II, III and IV may
03	themselves be prepared by standard techniques analogous
04	to those described above.
05 .	
06	It will be appreciated that a protected forms of a
07	peptide or analogue of the present invention are useful
08	novel intermediates and form an aspect of the

invention.

٥9 10 11

12

13

14

15

16

A peptide or analogue of the present invention may also be prepared on a solid phase support, for example a polyamide or a polystyrene resin, using amino acids . protected at the N-terminus, for example with the fluorenylmethyloxycarbonyl group or the t-butyloxycarbonyl group and with appropriate protection of any side-chain functional groups.

17 18 19

20

One such reaction scheme for solid-phase peptide synthesis is, for example, illustrated below.

02

ÒЗ

Solid Phase Scheme

A.A. = Amino acid t-BOC = t-Butyloxycarbonyl Fmoc = Fluorenylmethyloxy-

methyloxycarbonyl, i.e.

```
Resin
    Fmoc-A.A.(1) or
     t-BOC-A.A.(1)
Fmoc (or t-BOC) A.A.(1)-Resin
     Deprotection
H-A.A.(1)-Resin
     Coupling with
     Fmoc-A.A.(2) activated
Fmoc-A.A.(2)-A.A.(1)-Resin
     Deprotection
H-A.A.(2)-A.A.(1)-Resin
     Repeat coupling and
     deprotection steps until
     required peptide is
     obtained
Protected Peptide-Resin

↓ Resin cleavage and

    deprotection
Deprotected Peptide
```

- 17 -

This technique involves the addition of the first protected amino acid to a solid resin support. After removal of the protecting group (deprotection) the amino acid-resin is coupled with the next protected amino acid in its activated form. The deprotection/coupling procedures are repeated until the required peptide is obtained. The peptide is then cleaved from the resin before final removal of the protecting groups Alternatively, when desired or necessary, the protecting groups may be removed before cleavage of the peptide from the resin.

Advantageously the Fmoc group is the form of protection used for the α -amino function of the amino acids involved (but not for side chain protection).

However, the last amino acid in each synthesis is generally protected as its t-BOC or Fmoc derivative. This allows the peptide to remain fully protected on cleavage from the resin.

The use of alternative resins may also require the need for removal of protecting groups prior to resin cleavage. In this case it is likely that the Fmoc-protecting group would be used for $N\alpha$ protection throughout the syntheses.

The peptides and analogues of the present invention have smooth muscle relaxant activity such as gastro-intestinal, bronchodilator and vasodilator actions, and in addition, anti-ulcer activity. They may be useful in preventing the pain and constipation frequently encountered in some irritable bowel syndrome (IBS) patients and may be a useful new approach to duodenal ulcer therapy.

- 18 - 01 The present invention further provides a peptide or 02. analogue of the present invention, for use in a method 03 of treatment of the human or animal body. 04 05 Where the fragment or analogue is in the form of a salt 06 thereof, it should of course be understood that this is 07 a physiologically tolerable salt, which is 80 pharmaceutically acceptable. 09 10 The peptide or analogue of the invention may be 11 administered per se or, preferably, as a pharmaceutical 12 composition also including a pharmaceutically suitable 13 carrier. 14 15 Accordingly, the present invention provides a 16 pharmaceutical composition, which comprises a peptide 17 or analogue of the present invention, in admixture or 18 conjunction with a pharmaceutically acceptable carrier. 19 20 The preparation may, if desired, be in the form of a 21 pack accompanied by written or printed instructions for 22 23 use. 24 In accordance with conventional pharmaceutical practice 25 the carrier may comprise a diluent, filler, 26 disintegrant, wetting agent, lubricant, colourant, 27 flavourant or other conventional additive. 28 29 Preferably, a pharmaceutical composition of the 30 invention is in unit dosage form. 31 32 The suitable dosage range for compounds of the 33 invention may vary from compound to compound and may 34 depend on the condition to be treated. It will also 35 depend, inter alia, on the relation of potency to 36 absorbability and on the mode of administration chosen. 37

Suitable formulations are, for example, intravenous infusions, aerosols and enteric coated capsules.

The present invention further provides a method of treatment of a human or non-human animal, which comprises administering an effective, non-toxic, amount of a peptide or analogue of the present invention to a human or non-human animal; and a peptide or analogue of the present invention for use as a pharmaceutical, in particular for the treatment of disorders and complaints described below.

 A peptide or analogue of the present invention may be used to treat the following disorders and complaints; abnormalities of gut motility, e.g. hypermotility as in IBS or oesophageal spasm; peptic ulceration; bronchial spasm; vascular conditions such as hypertension and ischaemia; mental disorders.

 Conveniently, the active ingredient may be administered as a pharmaceutical composition hereinbefore defined, and this forms a particular aspect of the present invention.

A suitable dose is, for example, in the range of from 1 pg to 2.5 mg/kg i.v. in the rat. A possible daily dose for humans is, for example, 0.01 to 50 mg by intravenous infusion, 0.01 to 250 mg by aerosol or 0.1 to 500 mg by enteric coated capsule.

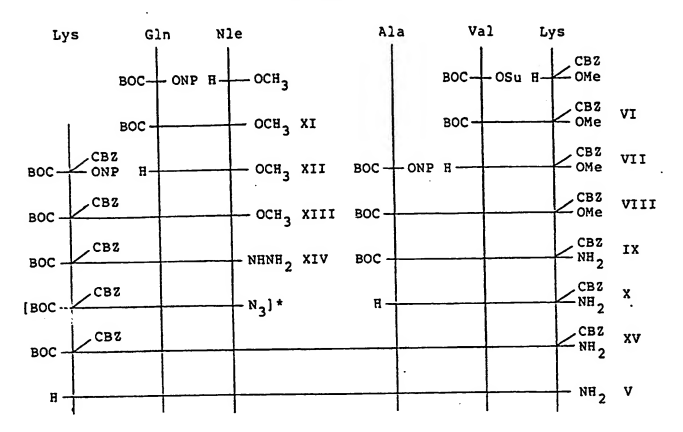
No adverse toxicological effects are indicated at the aforementioned dosage ranges.

In the following, the various derivatives protecting groups, reagents and solvents are referred to by abreviations for convenience.

01	- 20 -	0184309
02	Derivatives, Protecting	<u>Abbreviated</u>
03 🔆	Groups, Reagents, Solvents	<u>Designation</u>
04	Tertiary-butyl	But
05	Tertiary-butyloxycarbonyl	t-Boc
06	N-hydroxysuccinimide ester	OSu
07	Methyl ester	OMe
08	Trifluoroacetic acid	TFA
09	Dicyclohexylcarbodiimide	DCC
10	Benxyloxycarbonyl	€BZ
11	Dimethylformamide	DMF
12	Tetrahydrofuran	THF
13	p-Nitrophenyl ester	ONP
14	Hydrochloride salt	.HCl
15	Ethyl acetate	EtOAc
16	Methanol	MeOH
17	Ammonium Acetate	NH4OAC
18	1-Hydroxybenzotriazole	HOBT
19	Chloroform	CHC13
20	Pyridine	Pyr
21	n-Butanol	BuOH
22	Ammonium hydroxide	NH4OH
23	Sodium hydrogen carbonate	NaHCO3
24	Sodium chloride	NaCl
25	Ether	Et ₂ O
26	Sodium sulphate	Na 2504
27	Potassium hydroxide	КОН
28	Acetic acid	Acoh
29	•	· · · · · · · · · · · · · · · · · · ·
30	T.L.C. (Merck) silica gel plates) with	solvent systems
31	E_4 MeOH-CHCl ₃ (1 : 9)	
32		
33	H n-BuOH : AcOH : Pyr : H ₂ O (15 : 3 :	10:12)
34		
35	A_3 n-BuOH : AcOH : H_2 O (4 : 1 : 1)	
36		

01	- 21 -
02	Example 1
03	
04	H-Lys-Gln-Nle-Ala-Val-Lys-NH2 (V)
05	
06	The hexapeptide amide (V) was prepared as illustrated
07	in Scheme I and the experimental details are given
08	below.
09	

Scheme 1



* Prepared in situ.

t-Boc-Val-Lys(CBZ)-OMe (VI)

A mixture of t-Boc-Val-OSu (3.14 g, 10 mmol) and H-Mys(CBZ)-OCH3.HCl (3.31 g, 10 mmol) in THF (300 ml) was treated at room temperature with triethylamine (1.38 ml) and left stirring for 17 hr. The resulting solution was evaporated in vacuo and the residue was dissolved in EtOAc (500 ml). The organic solution was washed successively with water (2 x 200 ml), 5% citric acid (2 x 200 ml), water (2 x 200 ml) and dried over Na₂SO₄. The dried solution was filtered and evaporated in vacuo to give (VI) (4.9 g; 99%) as a foam. $R_f E_4 = 0.64.$

- 23 -

15 16

14

03 04

05

06 07

80

09

10 11

12 13

H-Val-Lys(CBZ)-OMe trifluoracetate (VII)

17 18

19

20

21 22

23

24

The protected dipeptide (VI) (4.9 g, 10 mmol) was dissolved in TFA (20 ml) and stirred for 10 minutes at room temperature. The solution was evaporated in vacuo, azeotroped with toluene (2 x 20 ml) and triturated with Et₂0 (2 x 50 ml). The mother liquors were decanted to leave the partially deprotected dipeptide as the trifluoroacetate salt VII (2.41 g; 51% $R_{f}E_{4} = 0.22.$

25 26 27

t-Boc-Ala-Val-Lys(CBZ)-OMe (VIII)

28 29

30 31

32

33 34

35

36

(VII) (2.4 g, 6.1 mmol) was added to t-Boc-Ala-ONP (1.9g, 6.1 mmol) in THF (50 ml) containing triethylamine (0.85 ml). The mixture was stirred at room temperature for 4 days, evaporated in vacuo and partitioned between EtOAc (250 ml) and water (100 ml). The organic layer was washed successively with 0.45M NH_4OH (4 x 50 ml), 2% citric acid (4 x 50 ml) and water (4 x 50 ml). The organic layer was dried over Na_2SO_4 ,

01	- 24 -
02	filtered and evaporated in vacuo. The residue was
03	recrystallised from EtOAc-hexane to give (VIII) (1.62
04	g; 47 %) as colourless microcrystals, mp 104°C.
05	$[\alpha]_D^{26} = -48^{\circ}C$ (C=1, MeOH).
06	
07	t-Boc-Ala-Val-Lys(CBZ)-NH2 (IX)
80	
09	(VII) (2.0 g, 35 mmol) was added to a solution of
10	ammonia (\underline{ca} . 50 ml) in methanol (50 ml). The mixture
11	was kept in a sealed pressure vessel for 24 hrs, then
12	evaporated to dryness. The residue was taken up in
13	EtOAc, evaporated to dryness and triturated with Et20
14	give (IX) (1.75 g; 90 %) as colourless microcrystals m
15	195-196°C $[\alpha]_D^{26} = -43.8^\circ$ (C=1, MeOH).
16	
17	H-Ala-Val-Lys(CBZ)-NH2.trifluoracetate (X)
18	
19	The protected tripeptide amide (IX) (1.0 g, 1.8 mmol)
20	was dissolved in cold TFA (10 ml). After 10 minutes,
21	the mixture was evaporated in vacuo and the residue was
22	triturated with ether (2 x 50 ml). The mother liquors
23	were decanted and the residue was dried under vacuum to
24	give (X) as a foam (0.85 g; 83 %).
25	
26	t-Boc-Gln-Nle-OMe (XI)
27	
28	t-Boc-Gln-ONP (12.3 g, 32 mmol) and HOBT (5.0 g, 37
29	mmol) were added to a solution of H-Nle-OMe.HCl (5.99
30	g, 33 mmol) and triethylamine (4.9 ml) in DMF (55 ml).
31	
32	The mixture was stirred at room temperature overnight,
33	EtOAc (100 ml) was added and the organic phase was
34	washed with 2 % citric acid, 0.45M NH4OH until free of
35	nitrophenol, 5 % NaHCO2, 2 % citric acid, water until

neutral, and a saturated solution of NaCl. The

- 25 -

 solution was dried over Na₂SO₄, filtered and concentrated in vacuo. Petroleum ether (bpt 40-60°C) was added, the precipitate was filtered, washed with the same solvent and dried in vacuo over silica gel to give (XI) (10.2 g, 85 %) mpt 108-109°C, [α D²⁶ = -14.89° (C=1, DMF), T.1.c. R_fA₃ = 0.72.

H-Gln-Nle-OMe.trifluoroacetate (XII)

The protected dipeptide ester (XI) (4.8 g, 13 mmol) was dissolved in cold TFA (40 ml). After 10 minutes, the TFA was removed in vacuo and dry ether (200 ml) was added. The ether was decanted and the residue was washed with more ether (100 ml). The oily material was dried over KOH to give (XII) as a white foam that was used immediately, T.1.c. $R_fA_3 = 0.42$.

BOC-Lys(CBZ)-Gln-Nle-OMe (XIII)

H-Gln-Nle-OMe.TFA salt (XII) (13 mmol) triethylamine (1.76 ml, 13 mmol), HOBT (2.16 g, 16 mmol) and BOC-Lys(CBZ)-ONP (7.6 g, 15 mmol) were dissolved in DMF (30 ml). The reaction mixture was kept basic with small amounts of triethylamine. The mixture was stirred overnight at room temperature, concentrated in vacuo and treated with unsymmetrical dimethylethylenediamine (2 equivs). After 2 hours, EtOAc was added and the product was isolated as described for compound (XI). The solid was washed with petroleum ether (bpt 40-60°C) and dried in vacuo over silica gel to give (XIII) $(5.5g; 65 \%) [\alpha_D^{26} = -16.49^{\circ} (C = 1, DMF) T.l.c. R_fA_3]$ 0.8 . 34

01	- 26 -
02	BOC-Lys(CBZ)-Gln-Nle-NHNH2 (XIV)
03	
04	The tripeptide methyl ester (XIII) (1 g, 1.6 mmol) was
05	suspended in DMF (5 ml), hydrazine (0.8 g, 0.78 ml, 16
06	mmol) was added and the mixture was stirred overnight.
07	The solvent was removed in vacuo and the residue
08	solidified using MeOH/EtOAc to give (XIV) (0.9 g, 90
09	%). T.1.c. $R_fA_3 = 0.70$ This was used immediately in
10	the next step.
11	
12	t-Boc-Lys(CBZ)-Gln-Nle-Ala-Val-Lys(CBZ) -NH2(XV)
13	
14	(XIV) (0.58 g, 0.9 mmol) was dissolved in anhydrous DMF
15	(18 ml) and cooled to -30°C. 4.56M HCl in dioxane
16	(0.90 ml) was added followed by t-butyl nitrite (0.12
17	ml). The reaction was left for 30 to 40 min at -30°C
18	then cooled to -60° C. Triethylamine (0.60 ml) was
19	added followed by the deprotected amide (X) (0.384 g,
20	0.68 mmol) and a further addition of triethylamine (0.1
21	ml). The mixture (M) was left to stand, reaching
22	ambient temperature over 2 days. A further amount of
23	(XIV) (0.29 g, 0.45 mmol) in DMF (10 ml) was treated at
24	-30° C with 4.5 M HCl in dioxane (0.45 ml), t-butyl
25	nitrite (0.06 ml) and triethylamine (0.3 ml) as
26	described above and added to the reaction mixture (M)
27	at -30° C. The whole was left to stand, reaching
28	ambient temperature over a further 4 days.
29	
30	The mixture was evaporated in vacuo and the residue was
31	triturated with EtOAc: MeOH (1:1) (50 ml) to give (XV)
32	(0.62 g; 85 %) as a greyish solid.

H-Lys-Gln-Nle-Ala-Val-Lys-NH2 (V)

The protected hexapeptide (XV) (0.2 g, 0.2 mmol) was dissolved in TFA (5 ml) and treated with HBr gas over 1 hr. The mixture was evaporated in vacuo and triturated with Et₂O (2 x 50 ml) to give (V) as a hydrobromide salt (0.13 g) R_fH = 0.14. This and a subsequent batch of product were purified by adsorption on to an ion exchange column (CM 25 Sephadex, Pharmacia) which was washed with 10-100 mmol NH₄OAc at pH 7. The product was eluted with 100 mmol NH₄OAc at pH 8.5.

Lyophilisation and subsequent preparative HPLC [µBondapak ODS.; CH₃CN: 50 mmol NH₄OAc(aq) (15: 85) gave (V) as an acetate salt (0.15 g) mp 253-255°C T.1.c. R_fH = 0.14, MH⁺ (FAB) = 685.

01	- 20 - •
02	Example 2
03	
04	H-Arg-Lys-Gln-Nle-Ala-Val-Lys-Lys-NH2.Acetate
05	(XVI)
06	
07	The octapeptide amide (XVI) was prepared as illustrated
08	in Scheme II and in the experimental details given
09	below.
10	. •
11	t-Boc-Lys(CBZ)-Lys(CBZ)-OCH3 (XVII)
12	
13	A mixture of t-Boc-Lys-(CBZ)-OH (1.14g, 3mmol),
14	Lys-(CBZ)-OCH3.HCl (0.99g, 3mmol), DCC (0.62g, 3mmol),
15	HOBT (0.41g, 3mmol) and triethylamine (0.42ml) in dry
16	amine-free DMF (20ml) was stirred for 17h. Work up as
17	described for VI gave XVII (1.2g; 61%) as colourless
18	microcrystals, mp 109-1100 (ex acetone-light petroleum
19	ether 40-60°) $[\alpha]_D^{26} = -11.9°$ (C=1 MeOH) $R_fE_4 = 0.71$.
20	
21	
22	H-Lys-(CBZ)-Lys-(CBZ)-OCH3.trifluoroacetate (XVIII)
23	
24	The protected dipeptide (XVII) (1.8g, 2.7mmol) was
25	partially deprotected as for VII to give XVIII as a
26	foam (1.8g; 99%) $R_fE_4 = 0.22$.
27	
28	
29	t-Boc-Val-Lys-(CBZ)-Lys-(CBZ)-OCH3 (XIX)
30	
31	A mixture of t-Boc-Val-OSu (0.86g 2.7mmol), XVIII
32	(1.8g, 2.7mmol) and triethylamine (0.4ml) in THF (50ml)
33	was stirred, under N_2 , for 2 days. Work up as
34	described for VI gave XIX (1.25g; 60%) as colourless
35	microcrystals, mp 145-1470 (ex EtoAc) $R_fE_4 = 0.40$
36	$[\alpha]_D^{26} = -24.59^{\circ}$ (C=1 MeOH).
37	

- 29 -01 H-Val-Lys-(CBZ)-Lys-(CBZ)-OCH3.trifluoroacetate (XX) 02 03 The protected tripeptide XIX (2.16g, 2.9mmol) was 04 partially deprotected as for VII to give XX as a flaky 05 solid (2.02g; 92%) 06 07 Boc-Ala-Val-Lys-(CBZ)-Lys-(CBZ)-OCH3 (XXI) 80 09 A mixture of XX (2.0g), Boc-Ala-ONP (1.0g, 3.2mmol), 10 HOBT (0.80g) and triethylamine (0.50ml) was stirred at 11 room temperature in DMF (5ml) for 24h. The mixture was 12 evaporated to & volume and taken up into CHCl3 13 The organic solution was washed successively 14 with 0.45M NH₄OH (4 x 50ml), 2% citric acid (4 x 50ml) .15 and water (4 x 50ml). The organic layer was dried over 16 Na₂SO₄, filtered and evaporated in vacuo to ½ volume. 17 The solution was chromatographed on Kieselgel 60 PF254 18 on a 'Chromatotron' and the product was eluted with an 19 increasing concentration of MeOH (0-5%) in CHCl3 to 20 give XXI (1.98g; 84%) as colourless microcrystals, mp 21 $175-177^{\circ}$ [α]_D²⁶ = -36.89 (C=1 MeOH). MH⁺ = 827 22 (FAB). 23 24 Boc-Ala-Val-Lys-(CBZ)-Lys-(CBZ)-NH2 (XXII) 25 26 XXI (1.78g, 2.2mmol) was added to a solution of ammonia 27 (ca. 50ml) in methanol (50ml). The mixture was kept in 28 a sealed vessel for 48h. The resulting precipitate was 29 filtered, washed with dry Et₂O to give XXII (1.78g; 30 98%) as colourless microcrystals mp 243-2440. 31 32 H-Ala-Val-Lys-(CBZ)-Lys-(CBZ)-NH2.trifluoroacetate 33 (XXIII) 34 35

The protected tetrapeptide XXII (1.75g, 2.2mmol) was

suspended in acetic acid (3ml), cooled to 100 and

36

treated with TFA (9ml). The solution was stirred for 20-25min. Work-up as described for VII gave XXIII (1.71g) $R_fE_1 = 0.7$.

H-Lys-(CBZ)-Gln-Nle-OMe (XXIV)

10 .

The protected tripeptide ester (XIII) (2.0g, 3mmol) was dissolved in cold TFA (12.6ml) and glacial acetic acid (5.4ml). After 25 minutes, the solvents were removed in vacuo and dry ether (100ml) was added. The ether was decanted and the residue was washed with more ether (100ml). The oily material was dried in vacuo over KOH to give the trifluoroacetate salt of (XXIV) as a white foam. The foam was dissolved in water (40ml) and a cold solution of sodium carbonate (0.15g) in water (10ml) added. The free base was extracted into ethyl acetate (100ml, then 4 x 30ml) and this organic phase was washed with water (2 x 20ml), saturated NaCl (20ml), dried over Na₂SO₄, filtered and evaporated in vacuo to give XXIV (1.5g; 89%). T.1.c. R_f = 0.3 in 30% MeOH/CHCl₃.

BOC-Arg(H⁺)-Lys-(CBZ)-Gln-Nle-OMe (XXV)

The free amine (XXIV) (1.5g, 2.8mmol) was dissolved in DMF (9ml). The solution was cooled, then BOC-Arg(H⁺)OH (1.23g, 4.5mmol), DCC (0.82g, 4mmol) and HOBT (0.57g, 4mmol) were added. After 2 hours, additional portions of BOC-Arg(H⁺)OH (0.45g, 1.6mmol), DCC (0.3g, 1.5mmol) and HOBT (0.2g, 1.6mmol) were added and the reaction was allowed to proceed for 3 days.

The dicyclohexylurea was removed by filtration and washed with DMF (3 x 5ml). The solvent was removed <u>in vacuo</u> and the residue was applied in methanol to a Sephadex LH-20 column (2.5 x 100cm) pre-equilibrated

with the same solvent. Fractions of 5ml were collected at a flow rate of lml/3mins. The fractions containing the desired product were pooled, evaporated and re-chromatographed under the same conditions.

The product was further purified on Kieselgel 60 Pf₂₅₄ using a 'Chromatotron' (20% MeOH/CHCl₃ as eluant) to give (XXV) (1.3g; 56%) $[\alpha]_D^{26} = -24.9^\circ$ (C = 1, MeOH).

BOC-Arg(H+)-Lys-(CBZ)-Gln-Nle-NHNH2 (XXVI)

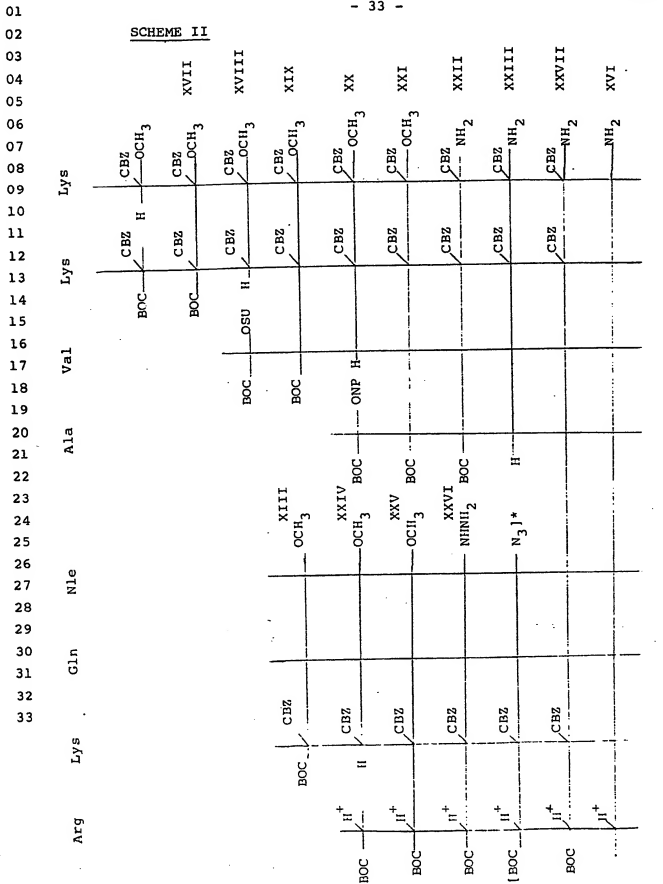
The tetrapeptide methyl ester (XXV) (1.3g, 1.6mmol) was suspended in methanol (6ml), hydrazine hydrate (0.8g, 0.78ml, 1.6mmol) was added, and the stirring was continued for 6 hours. The product was filtered, washed with cold methanol (3 x 10ml), water (8 x 5ml) and dried in vacuo to give (XXVI) (1.2g; 92%). The product was used immediately.

BOC-Arg(H+)-Lys-(CBZ)-Gln-Nle-Ala-Val-Lys-(CBZ)-Lys-(CBZ)-NH2.Chloride (XXVII)

The protected tetrapeptide (XXVI) (0.37g, 0.45mmol) was dissolved in anhydrous DMF (5ml) and cooled to -30°.

4.56M HCl in dioxane (0.45ml) was added followed by t-butylnitrite (0.06ml). The reaction was left for 30-40 min at -30° then cooled to -60°. Triethylamine (0.30ml) was added followed by the deprotected amide XXIII (0.28g, 0.3mmol) and a further addition of triethylamine (0.05ml). The mixture M2 was left to stand, reaching ambiant temperatures over 2 days. A further amount of XXVI (0.21g, 0.26mmol) in DMF (5ml) was treated at -30°C with 4.56M HCl in dioxane (0.25ml), t-butyl nitrite (0.04ml) and triethylamine (0.17ml) as described above and added to the reaction

01	- 32 -
02	mixture M_2 at -30°C. The whole was left to stand,
03	reaching ambiant temperature over a further 4 days.
04	
05	The mixture was treated with methanol and the whole
06	centrifuged. The resulting solid and mother liquors
07	were both shown to contain the desired product XXVII
08	$(1.25g) MH^+ = 1471 (FAB).$
09	H-Arg-Lys-Gln-Nle-Ala-Val-Lys-Lys-NH2.Acetate (XVI)
10	•
11	The protected octapeptide (XXVII) (1.25g, 0.85mmol) was
12	dissolved in TFA at 100 and treated with hydrogen
13	bromide gas for 2h. The whole mixture was evaporated
14	in vacuo, triturated with ether (4 x 15ml) and filtered
15	to give the free peptide as its hydrobromide salt
16	(1.0g). The peptide was purified with concomitant
17	conversion to an acetate salt, XVI (0.22g) (MH+ (FAB) =
18	969) in the same manner as that described for (V).



	0184;
01	- 34 -
02	Solid Phase Synthesised Peptides
03	
04	a)
05	The following examples were synthesised by solid phase
06	methods using the 4-hydroxymethylbenzoylnorleucyl
07	derivatised polydimethylacrylamide gel resin Pepsyn B
08	(1.0mequiv/g or 0.3mequiv/g) as supplied by Cambridge
09	Research Biochemicals Ltd.
10	
11 .	DMF was fractionally distilled in vacuo from ninhydrin
12	before use and stored over pre-activated molecular
13	sieves (4A). Piperidine was freshly distilled from a
14	suitable drying agent. Dichloromethane (A.R) was dried
15	over pre-activated molecular sieves (4A).
16	
17	The amino acids were chosen as their Fmoc-derivatives
18	with BCC- or t-Bu- side chain protection where
19	necessary.
20	
21	The symmetrical anhydride of the first amino acid
22	(2.5equiv), (prepared as described by E. Brown et al in
23	J.C.S. Perkin I, 1983, 80) was added to the resin (1
24	equiv) in DMF (10-15ml) in the presence of a catalytic
25	quantity of dimethylaminopyridine. The mixture was
26	agitated with N2 and the reaction was allowed to
27	proceed for lh. The resin was drained and the addition
28	procedure was repeated. The drained resin was then
29 :	washed with DMF (10-15ml x l min x 10).
30	The removal of the Fmoc protecting groups was achieved
31	by agitation of the peptide-resin with piperidine
32	(10ml; 20% in DMF) for 3 min then 7 min.

Subsequent addition of each amino acid was carried out using the Fmoc symmetrical amino acid anhydrides (2.5 equiv) or the preformed hydroxybenzotriazole ester (3.0 equiv) (from Fmoc-amino acid, DCC and HOBT).

 Amino acids containing amidic side chains (e.g. Gln or Asn) were coupled as their p-nitrophenyl activated esters (3.0equiv) in the presence of hydroxybenzotriazole (6.0equiv).

Fmoc-Arginine was coupled to the peptide resin via its hydroxybenzotriazole ester. This was prepared by suspending Fmoc-Arginine (10equiv) in DMF (10ml) and adding HOBT (30 equiv). The clear solution was added to the resin and agitated for 1 minute. DCC (10 equiv) was then added and the reaction was allowed to proceed to completion.

The final amino acid in the chosen sequence was added as its $N\alpha$ Boc derivative either as the symmetrical anhydride or as the preformed hydroxybenzotriazole ester.

Boc-Arginine was coupled as its hydrochloride and activated by addition of DCC (5 equiv) to the protected hydrochloride salt (10 equiv) in DMF (10-15ml) 5 minutes prior to addition of the whole reaction mixture to the peptide-resin (1 equiv).

In some cases, Fmoc-amino acid anhydrides (eg Phe, Ala, Gly) coprecipated with DCU during their formation. In these cases, the anhydrides were prepared in the presence of 10% DMF in dichloromethane.

Dichloromethane was removed in vacuo before addition of the whole mixture to the peptide resin.

Couplings in general were carried out for 1-2h and repeated if necessary. Completeness of acylation was verified by a qualitative Kaiser ninhydrin test as described by E. Kaiser et al in Anal. Biochem. (1970) p.34.

0184309

The vessel

- 36 -

01

Peptide cleavage from the resin was accomplished via 02 ammonolysis to provide the protected peptide amide. 03 this end, when the final coupling was complete, the 04 peptide-resin was washed with DMF (10-15ml x 1 min x 05 10), anhydrous dichloromethane (10-15ml x 1 min x 10) 06 and dry ether (10ml x l min x 10). The collapsed resin 07 was dried over silica gel for l hour in a vacuum 80 desiccator: The resin was re-swollen as previously 09 described, drained and treated with a saturated 10 solution of ammonia in methanol at -100. 11 was sealed and allowed to reach ambiant temperatures 12 for 2 days. The apparatus was cooled, opened and the 13 contents were allowed to warm to room temperature. 14 15 suspension was filtered under suction and the resulting 16 residue was washed with methanol (5 x 5ml) and DMF (5 x17 The combined washings and filtrate were evaporated in vacuo. 18 The resulting residue was triturated with dry ether and filtered to give the 19

protected peptide.

20 21 22

23

24 25

26

27

28

29

30

31

32

The final acidolytic deprotection procedure removed all protecting groups (e.g. BOC, t-Bu) from the peptide amide. Thus the protected peptide was dissolved in trifluoroacetic acid (4ml/100mg of peptide) and stirred at room temperature for 3h. In some cases, hydrogen bromide gas was bubbled though the mixture during this The mixture was evaporated in vacuo and the resulting solid was triturated with dry ether (7 x 5ml) to give the required peptide either as its trifluoroacetate or its hydrobromide salt. The peptides were purified by one or a combination of methods listed below.

33 34

01	- 37 -
02	(a) Conversion to acetate salt.
03	
04	The peptide salt was dissolved in a minimum amount of
05	water and passed down a strong anion exchange resin
06	which was in its acetate form (e.g. Sephadex
07	QAE-A-25). Eluant was fractionated and the fractions
08	containing desired materials were lyophilised.
09	
10	b) Selective adsorbtion chromatography
11	
12	The peptide salt was dissolved in a minimum amount of
13	water and adsorbed onto a weak cation exchange resin
14	(e.g. Sephadex CM-25). The peptide acetate was
15	recovered during elution with an increasing
16	concentration of NH_4OAc (0.05M - 0.5M) at pH 7, an
17	increasing pH gradient (pH 7 - pH 9) or a combination
18	of both.
19	
20	c) High Performance Liquid Chromatography. HPLC
21	·
22	The peptide was purified by preparative HPLC on
23	reverse phase C_{18} silica columns (e.g. μ bondapak,
24	Hypersil ODS).
25	
26	The peptides were characterised by 24h acidolytic
27	cleavage and PITC derivatised amino acid analysis
28 :	(Waters Picotag system) and fast atom bombardment (FAB
29	mass spectrometry (Jeol DX 303).
30	Example 3
31	
32	H-Leu-Arg-Lys-Gln-Nle-Ala-Val-Lys-Lys-NH2.Acetate
33	(XXVIII)
34	
35	XXVIII was prepared using the 0.3mequiv/g Pepsyn B
36	resin.
37	$[MH]^+ = 1081 (FAB).$

01	- 38 -
02	Amino acid analysis. Glu (1.0) Arg (1.0) Ala (1.0) Val
03	(0.88) Nle (1.0) Leu (1.0) Lys (2.88).
04	
05	
06	Example 4
07	
08	H-Lys-Gln-Nle-Ala-Leu-Lys-NH2.Acetate (XXIX)
09	
10	XXIX was prepared using the 1.0 mequiv/g Pepsyn B
11	resin. [MH]+ (FAB) = 699. Amino acid analysis: Glu
12	(0.9), Ala (0.8), Leu (0.8), Nle (0.8), Lys (1.8).
13	
14	
15	Example 5
16	
17	H-Lys-Gln-Nle-Ala-Val-Orn-NH2.Acetate (XXX)
18	
19	XXX was prepared using the 1.0mequiv/g Pepsyn B resin
20	[MH] $^+$ (FAB) = 671. Amino acid analysis; Glu (1.01),
21	Ala (0.9), Val (0.8), Nle (1.1), Lys (1.0), Orn (1.1).
22	
23	
24	Example 6
25	
26	H-Orn-Gln-Nle-Ala-Val-Orn-NH2.Acetate (XXXI)
27	
28	XXXI was prepared using the 1.0mequiv/g Pepsyn B resin
29	$[MH]^+$ (FAB) = 657.
30	
31	Example 7
32	
33	H-Lys-Gln-Leu-Ala-Val-Lys-NH2.Acetate (XXXII)
34	
35	XXXII was prepared using the 1.0mequiv/g Pepsyn B resin
36	$[MH]^+$ (FAB) = 685.
37	•
20	

01	- 39 -
02	Example 8
03	
04	H-Arg-Lys-Gln-Nle-Ala-Val-Lys-Lys-Tyr-Leu-NH2.Acetate
05	(XXXIII)
06	
07	XXXIII was prepared using the 0.3mequiv/g Pepsyn B
08	resin. $[MH]^+$ (FAB) = 1245.
09	
10	
11	Example 9
12	•
13	H-Lys-Gln-Nle-Ala-Val-Lys-Lys-Tyr-Leu-NH2.Acetate
14	(XXXIV)
15	
16	XXXIV was prepared using the 0.3mequiv/g Pepsyn B
17	resin. $[MH]^+$ (FAB) = 1089.
18	
19	b) Use of the Beckmann model 990B Peptide
20	Synthesiser
21	
22	The following example was synthesised using leucine
23	resin ester. This was prepared by reacting
24	chloromethylated resin (3.5g, 0.7mequiv Clg^{-1} ; 1%
25	cross-linked styrene/divinylbenzene as supplied by
26	Merseyside Laboratories) at 50°C, for 17h, in DMF
27	(40ml) with the anhydrous cesium salt obtained from
28	Boc-L-leucine monohydrate (0.5g, 2mmol). The resulting
29 .	Boc-L-leucine resin ester was exhaustively washed with
30	DMF, 50% aqueous DMF, H2O, DMF and finally CH2Cl2, then
31	dried (3.68g; 0.22mmol leucine/g).
32	
33	Removal of the BOC group (from 3.6g resin) was achieved
34	by reaction with 50% TFA in CH_2Cl_2 (50ml) for 5 min
35	then 25 min. The leucine resin ester salt was washed
26	with Cucle (7 v 50ml) noutralised with 10%

01 .	- 40 -
02	di-isopropylamine in CH_2Cl_2 for 5 min (3 x 50ml) and
03	washed with CH_2Cl_2 (8 x 50ml).
04	·
05	The first amino acid of the required sequence was
06	coupled to the leucine resin ester by the following
07	procedure. Fmoc-L-tyrosine-t-butyl ether (6mmol) and
08	di-isopropylcarbodiimole (6mmol) were reacted with the
09	leucine resin ester in CH_2Cl_2/DMF (35ml) for 12h then
10	the $Fmoc$ -Tyr(Bu ^t)-Leu-resin ester was washed with
11	CH_2Cl_2 (5 x 50ml).
12	
13	To remove the Fmoc protecting group, the peptide resin
14	was washed with DMF (5 x 50ml), reacted with 55% \cdot
15	piperidine in DMF (50ml) for 5 min then 20 min, then
16	washed with DMF (5 x 50ml).
17	
18	Subsequent Fmoc amino acids were coupled using the
19	procedure described above except for Fmoc-glutamine
20	which was incorporated using the HOBT/DCC
21	pre-activation procedure of König and Geiger (Chem.
22	Ber., 103, 788-98, (1970)).
23	
24	Couplings, in general, were carried out for 1-2h and
25	repeated if necessary. Completeness of acylation was
26	verified by a qualitative ninhydrin test as described
27	by E. Kaiser et al, in Anal. Biochem., (1970), p34.
28	
29	Deprotection was carried out by reaction with 55%
30	piperidine in DMF, as described above, followed by
31	reaction with a mixture of TFA (45ml), CH2Cl2 (45ml),
32	anisole (10ml) and methionine (1g) for 84 min. The
33	peptide resin was then washed with CH_2Cl_2 (5 x 50ml)
34	and dried.
35	

01	₹♣
02	Peptide cleavage from the resin was accomplished via
03	ammonolysis to provide the peptide amide. To this end
04	the peptide resin was stirred with ammonia-saturated
05	methanol (120ml) for 44h, filtered and washed with
06	methanol. Evaporation of the combined washings
07	followed by lyophilisation from aqueous acetic acid
08	gave the crude peptide amide. The ammonolysis was
09	repeated if FAB mass spec showed the presence of
10	peptide ester.
11	
12	Purification was carried out via HPLC on a Lichoprep
13	RPS column (25 x 1.6cm) with 0.1% aqueous TFA (A) and
14	90% acetonitrile/10% A (B) as a gradient from 0% B to
15	100% B over 60 min at 12ml/min.
16	
17	Example 10
18	
19	H-Lys-Gln-Met-Ala-Val-Lys-Lys-Tyr-Leu-NH2. Acetate
20	(VXXXV)
21	
22	$[MH]^+$ (FAB) = 1007
23	
24	Amino acid analysis; Glu (0.92), Ala (0.92), Tyr
25	(0.98), Val (0.99), Met (0.85), Leu (1.08), Lys (3.24)
26	
27	Example 11 and 12
28	~ ·
29	The following examples are prepared in accordance with
30	the methods described for examples 3 to 9.
31	
32	H-Lys-Gln-Ala-Val-Lys-NH2 (XXVI)
33	
34	H+
35	
36	H-Leu-Arg-Lys-Gln-Nle-Ala-Val-Lys-NH2 (XXVII)

01	- 42 -
02	Pharmacological Data.
03	I Colonic Motility
04	
05	(a) <u>In vivo</u>
06	
07	Male albino rats, Wistar strain (Charles River UK)
08	300-500 g were anaesthetised with urethane. A segment
09	of proximal colon was prepared for intraluminal
10	pressure recording after the method of Maggi and Meli
11	(Maggi, C.A. and Meli, A. (1982), J Pharmacol. Methods
12	8, 39-46). The activity of the compound (V) was
13	assessed from its action on the spontaneous motility of
14	the preparation after intravenous administration. The
15	compound was found to be active at 3 µmol/kg.
16	
17	(b) <u>In vitro</u>
18	
19	Segments of circular muscle cut from the proximal color
20	of rats were mounted in Krebs solution in isolated
21	tissue baths after the methods of Couture et al.
22	(Couture R. et al. (1981), Can. J. Physiol.
23	Pharmacol., 59, 957-964; and Couture R. et al., (1982),
24	Pharmacol., 24, 230-242). The activity of the compound
25	V was assessed from the effect on the spontaneous
26	contractile responses generated by this tissue.
27	The ED ₅₀ was found to be 10^{-5} M.

- 43 -

II Anti-ulcer Activity

03 04

05

06

07

.08

01

02

Anti-ulcer activity may be related to the enhanced capacity to dispose of gastric acid. Acid disposal capacity may be enhanced by increased intestinal secretions and enhanced acid disposal capacity is believed to be useful in the treatment of peptic ulcer disease.

09 10

11

Method for estimating the acid disposal capacity of the rat proximal duodenum

12 13 14

15

16

17

18

19

20

21

22

23

24

25

26

27

28

29

30

31

Male Wistar rats, 180-250g bodyweight, fasted overnight, are anaesthetized with urethane (150mg/100g bodyweight i.m.). The trachea is cannulated, and a gastric cannula, 0.5cm i.d., 3cm long, is inserted into the non-glandular forestomach via a mid-line abdominal incision. The intragastric cannula is exteriorized via a stab wound in the body wall. A triple lumen catheter, 0.3cm o.d., is passed via the gastric cannula through the pylorus. The duodenum is ligated 1cm below the pylorus, and the pylorus ligated around the cannula, thus creating a lcm proximal duodenal pouch excluding pancreatic and biliary secretions. The two ligatures enclosing the duodenal pouch are placed so as to avoid occluding the blood supply to the duodenal segment. Gastric secretions are allowed to drain freely from the gastric cannula. Compounds are administered dissolved in 0.9% sodium chloride (saline) as a 1.2 ml/h infusion via a catheter inserted in a jugular vein.

32 33 34

35

36

The triple lumen catheter is connected as follows. Lumen 1 delivers perfusing medium at 0.1ml/min via a peristaltic pump. 01 - 44 -

Lumen 2 collects the perfusate and delivers it to a flow cell containing a pH microelectrode. Outflow pH is recorded throughout the experiment. Lumen 3 is connected to a pressure transducer to monitor intraluminal pouch pressure. Body temperature is maintained at 34°C throughout the experiments.

- 14

After preparation, the duodenal segment is perfused with saline, adjusted to pH 6.5 with hydrochloric acid, for 30 minutes. The perfusing medium is then changed successively to saline adjusted with hydrochloric acid to pH 4, 3.5, 3 and 2.5 in increasing order of acidity. Each solution is perfused for 40 minutes. At the end of the pH 2.5 infusion period, saline pH 6.5 is perfused for 30 minutes, and the descending pH series repeated. This procedure produces two series of input pH/output pH values, designated 1st and 2nd passes.

A group size of 6 animals or larger is used and the effect of compounds on the output pH compared to control data determined. For comparisons between groups, Student's 't' test is used. Significance is taken at P<0.05.

 The compound of example 4 caused a significant increase in acid disposal at input pH 3 and 2.5 on the first pass and input pH 2.5 on the second pass at a dose of 150nmol/kg/h, and at input pH 2.5 on the first pass at a dose of 30nmol/kg/h.

U	T	
0	2	

Claims

C

1. A peptide comprising, in sequence, units
selected from the amino acid residues 11 to 23 of VIP
and consisting at least of the amino acid residues 15
to 20, or an analogue thereof wherein one or more of
the amino acid residues is replaced by an equivalent
other amino acid, or a pharmaceutically acceptable salt
thereof.

2. A peptide according to claim 1 wherein the amino acid units are selected from residues 13 to 23 or 11 to 21 of VIP or an analogue thereof as defined in claim 1.

3. A peptide according to claim 1 or 2 wherein all the amino acid units are in the L-form.

4. An analogue of a peptide according to any one of claims 1 to 3 in which one or more of the amino acid residues 15 to 20 is replaced by an equivalent other amino acid and any additional amino acid residues present correspond to those in VIP.

5. A peptide or analogue according to any one of claims 1 to 4 wherein the carboxy - terminus of the peptide or analogue is in the form of the unsubstituted amide.

6. A peptide or analogue according to any one of claims 1 to 5 wherein the amino-terminus of the peptide is in the form of the unsubstituted amine.

01	- 2 -
02	 A polypeptide of the general formula
03	
04	$X - (Y')_n - Y_{15}Y_{16}Y_{17}Y_{18}Y_{19}Y_{20} - (Y'')_m - Z$
05	
06	in which
07	
08	X represents a hydrogen atom or an amine-protecting
09	group, preferably a hydrogen atom;
10	
11	(Y') _n represents a direct bond or
12	
13	Y ₁₁ Y ₁₂ Y ₁₃ Y ₁₄ , Y ₁₂ Y ₁₃ Y ₁₄ , Y ₁₃ Y ₁₄ or Y ₁₄
14	
15	in which
1 6	
17	Y_{11} represents Thr, Ser or the residue of another
18	hydroxy amino acid,
19	·
20	$ m Y_{12}$ represents Arg or the residue of another basic
21	amino acid,
22	
23	Y_{13} represents Leu, Phe or the residue of another
24	hydrophobic amino acid,
25	
26	Y_{14} represents Arg or the residue of another basic
27	amino acid;
28	
29	Y_{15} represents Lys or the residue of another basic
30	amino acid, e.g. Orn,
31	
32	Y ₁₆ represents Gln or the residue of another carbox-
33	amido amino acid,
34	
35	Y ₁₉ represents Val or the residue of another hydro-
36	phobic amino acid,
37	· •

01	- 3 -
02	Y20 represents Lys or the residue of another basic
03	amino acid, e.g. Orn,
04	
05	(Y'') _m represents a direct bond or
06	
07	Y ₂₁ , Y ₂₁ Y ₂₂ or Y ₂₁ Y ₂₂ Y ₂₃
08	·
09	in which
10	
11	Y21 represents Lys or the residue of another basic
12	amino acid,
13 .	•
14	Y22 represents Tyr or the residue of another
15	hydrophobic amino acid,
16	·
17	Y23 represents Leu or the residue of another hydrophic
18	amino acid; and
19	
20	Z represents a hydroxyl group, or a group of the
21	formula OR such that COZ represents an ester, or a
22	hydrazino group such that COZ represents a hydrazide,
23	or NH ₂ such that COZ represents an amide; and
24	pharmaceutically acceptable salts thereof.
25	
26	8. The hexapeptide
27	
28	Lys Gln Y Ala Val Lys
29	
30	or Lys Gln Y ala Leu Lys
31	
32	wherein Y represents Met or Nle.
33	
34	9.
35	
36	H-Lys-Gln-Nle-Ala-Val-Lys-NH2,
37	

01	- 4 -
02	H-Arg-Lys-Gln-Nle-Ala-Val-Lys-Lys-NH2,
03	·
04	H-Leu-Arg-Lys-Gln-Nle-Ala-Val-Lys-Lys-NH2,
05	
06	H-Lys-Gln-Nle-Ala-Leu-Lys-NH2,
07	·
08	H-Lys-Gln-Nle-Ala-Val-Orn-NH2,
09	·
10	H-Orn-Gln-Nle-Ala-Val-Orn-NH2,
11	
12	H-Lys-Gln-Leu-Ala-Val-Lys-NH2,
13	
14	H-Arg-Lys-Gln-Nle-Ala-Val-Lys-Lys-Tyr-Leu-NH2,
15	
16	H-Lys-Gln-Nle-Ala-Val-Lys-Lys-Tyr-Leu-NH2,
17	
18	H-Lys-Gln-Met-Ala-Val-Lys-Lys-Tyr-Leu-NH2,
19	
20	H-Lys-Gln-Nle-Ala-Val-Lys-Lys-NH2 or
21	H+
22	
23	H-Leu-Arg-Lys-Gln-Nle-Ala-Val-Lys-NH2, or a
24	pharmaceutically acceptable salt of any of the
25	foregoing.
26	
27	10. A compound according to any one of claims 1 to 9
28	in pharmaceutically acceptable form.
29	
30	11. A compound according to any one of claims 1 to
31	10 whenever prepared synthetically.
32	
33	12. A process for the preparation of a peptide or
34	analogue according to any one of claims 1 to 11, which
35	comprises coupling a suitable amino acid or amino acid
36	sequence in which the carboxyl group is activated with
37	an appropriate amino acid or amino acid sequence and

repeating, if necessary, the coupling procedure until there is obtained a peptide comprising, in sequence, units selected from the amino acid residues 11 to 23 of VIP consisting at least of the amino acid residues 15 to 20, or an analogue thereof in which one or more of the amino acid residues is replaced by an equivalent other amino acid, wherein, if desired or required, non-reacting functional groups are protected during the coupling procedure and, if desired, subsequently deprotected, and optionally thereafter forming a pharmaceutically acceptable salt thereof.

- 13. A pharmaceutical composition comprising a peptide or analogue according to any one of claims 1 to 11 or a pharmaceutically acceptable salt thereof and a pharmaceutically acceptable carrier.
- 14. A peptide or analogue according to any one of claims 1 to 11 or a pharmaceutically acceptable salt thereof for the use as a pharmaceutical.
- 15. A peptide or analogue according to any one of claims 1 to 11 or a pharmaceutically acceptable salt thereof for use in the treatment and/or prophylaxis of ulcers.